Assignment 04

PUBH 8878

Requirements

- Show complete mathematical work for any derivations.
- Submit well-documented R code with clear comments and a fixed seed (set.seed(8878)).
- Interpret results in a **genetic/biological** context where applicable.
- Submit the rendered PDF only (do not submit the source .qmd).

A helpful vignette on using cmdstanr is available at https://mc-stan.org/cmdstanr/articles/cmdstanr.html.

Problem 1: Population Substructure and Allele Frequency Estimation (30 pts)

Let there be n diploid individuals sampled at random from a population made of subpopulations $k=1,\ldots,K$. In subpopulation k, the allele A frequency is p_k , and let w_k be the fraction of sampled individuals from subpopulation k ($\sum_k w_k = 1$). The "global" allele frequency we want to estimate is the mixture average

$$p = \sum_{k} w_k p_k.$$

Let n_{AA} , n_{Aa} , n_{aa} be the counts of genotypes AA, Aa, aa in the sample of size n. The standard estimator of the allele frequency is

$$\hat{p} = \frac{2n_{AA} + n_{Aa}}{2n}.$$

(a) Show that in the presence of population substructure, \hat{p} is unbiased.

Hint: Let D_i be the A-dosage for individual i $(D_i \in \{0,1,2\})$. Under HWE within k, $D_i \mid Z_i = k \sim \text{Bin}(2, p_k)$. Use LOTUS: $\mathbb{E}[D_i] = \sum_k w_k \mathbb{E}[D_i \mid Z_i = k]$.

(b) What is $Var(\hat{p})$ under random mixture sampling (i.i.d. individuals from the mixture)? Assume each subpopulation is in HWE. Compare to the case with no substructure (K = 1).

Hint: Use the law of total variance on D_i :

 $\mathrm{Var}(D_i) \,=\, \mathbb{E}\{\mathrm{Var}(D_i \mid Z)\} \,+\, \mathrm{Var}\{\mathbb{E}(D_i \mid Z)\}, \text{ with } \mathrm{Var}(D_i \mid Z = k) \,=\, 2p_k(1-p_k) \text{ and } \mathbb{E}(D_i \mid Z)\}$ $\mathbb{E}(D_i \mid Z=k) = 2p_k.$

Define

$$\bar{p} = \sum_{k} w_k p_k \tag{1}$$

$$\bar{p} = \sum_k w_k p_k \tag{1} \label{eq:power}$$

$$\operatorname{Var}_w(p_k) = \sum_k w_k (p_k - \bar{p})^2, \tag{2} \label{eq:power}$$

and note $\mathbb{E}[p_k(1-p_k)] = \bar{p}(1-\bar{p}) - \operatorname{Var}_w(p_k)$.

(c) Now consider a stratified sample: take exactly n_k individuals from subpopulation k $(\sum_k n_k = n; \text{ write } w_k = n_k/n)$. Maintain HWE within subpopulations. Derive $\text{Var}(\hat{p})$ under this design and compare it to your answer in (b). State which design yields the larger variance, and by how much, in terms of $Var_w(p_k)$.

Hint: Write $\hat{p} = \sum_k w_k \hat{p}_k$ with $\hat{p}_k = \frac{1}{2n_k} \sum_{i:Z_i=k} D_i$. Use independence across strata and $\operatorname{Var}(\hat{p}_k) = \frac{p_k(1-p_k)}{2n_k}.$

Problem 2: Population Substructure, LD, and Association Testing (40 pts)

Let X be the genotype dosage (0/1/2) copies of the effect allele) at a tag SNP and X_c at a causal SNP. The causal SNP has effect size β_c on quantitative trait Y. The observed effect from simple regression of Y on X is β_{obs} .

Convenience (scaling): Work with standardized genotypes

$$\tilde{X} = \frac{X - \mathbb{E}[X]}{\sqrt{\mathrm{Var}(X)}}, \qquad \tilde{X}_c = \frac{X_c - \mathbb{E}[X_c]}{\sqrt{\mathrm{Var}(X_c)}}.$$

With this scaling, $\beta_{\rm obs}=r\,\beta_c$ exactly, where $r={\rm Corr}(\tilde{X},\tilde{X}_c)={\rm Corr}(X,X_c).$

Assume individuals are sampled i.i.d. from a mixture with $\Pr(Z=k)=w_k,\;\sum_k w_k=1.$ Within each subpopulation k: - HWE holds at each locus, - LE (no within-k LD) holds between X and X_c .

Define

$$\bar{p} = \sum_{k} w_k p_k, \quad \bar{p}_c = \sum_{k} w_k p_{c,k} \tag{3}$$

$$\operatorname{Var}_w(p_k) = \sum_k w_k (p_k - \bar{p})^2 \tag{4}$$

$$\mathrm{Cov}_w(p_k,p_{c,k}) = \sum_k w_k(p_k - \bar{p})(p_{c,k} - \bar{p}_c) \tag{5} \label{eq:5}$$

Part A (10 pts): Correlation induced by population structure

Let the allele frequencies at the tag and causal SNPs be p_k and $p_{c,k}$ in subpopulation k. Show that

$$r = \frac{\operatorname{Cov}(X, X_c)}{\sqrt{\operatorname{Var}(X)\operatorname{Var}(X_c)}},$$

and express $Cov(X, X_c)$, Var(X), and $Var(X_c)$ in terms of $\{w_k, p_k, p_{c,k}\}$.

Hint: Law of total covariance:

 $\text{Cov}(X,X_c) = \mathbb{E}[\text{Cov}(X,X_c \mid Z)] + \text{Cov}(\mathbb{E}[X \mid Z],\mathbb{E}[X_c \mid Z]). \text{ Under LE, the first term is 0.}$ Use $\mathbb{E}[X \mid Z = k] = 2p_k$.

Part B (10 pts): Bias from ignoring structure in the trait

Suppose population structure also affects the trait mean: $\mathbb{E}[Y \mid Z = k] = \mu_k$. Consider the model $Y = \mu_Z + \beta_c X_c + \varepsilon$ with $\mathbb{E}[\varepsilon] = 0$. Show that the naïve regression of Y on X (without structure covariates) is biased:

$$\hat{\beta}_{\text{na\"{i}ve}} \approx \frac{\beta_c \operatorname{Cov}(X, X_c) + \operatorname{Cov}(X, \mu_Z)}{\operatorname{Var}(X)} = r \, \beta_c + \underbrace{\frac{\operatorname{Cov}(X, \mu_Z)}{\operatorname{Var}(X)}}_{\text{bias}}.$$

Under the assumptions above, prove that $\mathrm{Cov}(X,\mu_Z)=2\,\mathrm{Cov}_w(p_k,\mu_k)$, and give the bias in terms of w_k,p_k,μ_k .

 $\mathit{Hint:}\ \mathrm{Cov}(X,\mu_Z) = \mathrm{Cov}(\mathbb{E}[X\mid Z],\mu_Z) = \mathrm{Cov}(2p_Z,\mu_Z).$

Part C (20 pts): Brief interpretation

In a few sentences each:

1. Explain why $r \neq 0$ can arise even if **within** each subpopulation there is no LD. What feature of the mixture induces it?

- 2. Give a sign-consistent example: if subpopulations with larger p_k also have larger μ_k , what is the expected direction of the naïve bias?
- 3. Name **two** standard strategies to mitigate both components of bias (structure-induced r and trait mean differences) in practice.

Problem 3: Bayesian Analysis (30 pts)

Part A (10 pts): Beta-Binomial conjugacy

- 1. With prior Beta(4, 18) and data x = 11 successes out of n = 27, write the posterior distribution for p.
- 2. Compute the posterior mean and a **central 95% credible interval** in R using **qbeta**. Compare to the MLE $\hat{p} = 11/27$. Briefly interpret the *shrinkage*.
- 3. **Sensitivity:** repeat with priors Beta(1,1) and Beta(8,32). Summarize how the posterior mean and width change across priors, and why.

Part B (10 pts): Beta-Binomial in Stan

- 1. Write a Stan model to estimate the allele frequency p from Binomial data with a Beta(4,18) prior. Use x=11, n=27.
- 2. Run the model in R using cmdstanr. Check convergence and effective sample size; report \hat{R} and bulk ESS for p.
- 3. **Summarize** the posterior mean and a central 95% credible interval. Compare to your analytical result from Part A.

You will need to install cmdstanr and cmdstan if you haven't already. Please follow installation instructions at https://mc-stan.org/cmdstanr/

Boilerplate is provided below. You will need to set eval to TRUE to run the code when knitting the final document:

```
library(cmdstanr)

stan_beta_binomial <- "
data {
  int<lower=0> n;
  int<lower=0, upper=n> x;
```

```
real<lower=0> a;
  real<lower=0> b;
parameters {
real<lower=0, upper=1> p;
}
model {
 // TODO: prior on p
 // Example: p ~ beta(a, b);
  // TODO: likelihood
 // Example: x ~ binomial(n, p);
generated quantities {
real logit_p = logit(p);
 int x_rep = binomial_rng(n, p);
library(cmdstanr)
set.seed(8878)
writeLines(stan_beta_binomial, con = "beta_binomial.stan")
mod_bb <- cmdstan_model("beta_binomial.stan")</pre>
fit_bb <- mod_bb$sample(</pre>
  data = list(n = 27, x = 11, a = 4, b = 18),
  seed = 8878, chains = 4, parallel_chains = 4,
  iter_warmup = 1000, iter_sampling = 1000
# TODO: check convergence and summarize posterior
```

Part C (10 pts): ABO blood group frequencies in Stan (missing AB phenotype)

We observe phenotype counts in a population sample where **AB** individuals are not sampled:

- $n_A = 725$
- $n_B = 258$
- $n_O = 1073$

Under HWE with allele frequencies $\mathbf{p}=(p_A,p_B,p_O)$, the **unconditional** phenotype probabilities are

$$\Pr(A) = p_A^2 + 2p_A p_O, \quad \Pr(B) = p_B^2 + 2p_B p_O, \quad \Pr(AB) = 2p_A p_B, \quad \Pr(O) = p_O^2.$$

Because AB is missing, the **observed** category probabilities are the renormalized values

$$q_A = \frac{\Pr(A)}{1 - \Pr(AB)}, \quad q_B = \frac{\Pr(B)}{1 - \Pr(AB)}, \quad q_O = \frac{\Pr(O)}{1 - \Pr(AB)}.$$

- 1. Write a Stan model that estimates (p_A, p_B, p_O) with prior Dirichlet(1, 1, 1) and a Multinomial likelihood on (n_A, n_B, n_O) using (q_A, q_B, q_O) .
- 2. Run the model and check convergence (report \hat{R} and ESS).
- 3. **Prior sensitivity:** re-run with $\alpha = k (0.26, 0.09, 0.65)$ for $k \in \{1, 10, 100\}$. Summarize how posterior means and credible intervals change with k, and why.

Boilerplate is provided below:

```
stan_abo_missing_ab <- "
data {
  int<lower=0> n_A;
  int<lower=0> n_B;
  int<lower=0> n_0;
  vector<lower=0>[3] alpha;
transformed data {
  int N = n_A + n_B + n_0;
  int y[3] = \{ n_A, n_B, n_0 \};
parameters {
  simplex[3] p;
transformed parameters {
  // Unconditional phenotype probabilities under HWE:
 real PrA = square(p[1]) + 2 * p[1] * p[3];
  real PrB = square(p[2]) + 2 * p[2] * p[3];
  real PrAB = 2 * p[1] * p[2];
  real Pr0 = square(p[3]);
  // Observed (AB excluded): renormalize by (1 - PrAB)
  simplex[3] q;
```

```
real denom = 1 - PrAB;
    q[1] = PrA / denom;
   q[2] = PrB / denom;
   q[3] = Pr0 / denom;
  }
model {
 // TODO: prior on allele frequencies
 // Example: p ~ dirichlet(alpha);
 // TODO: likelihood for observed counts
 // Example: y ~ multinomial(q);
generated quantities {
 // Unconditional phenotype probabilities (optional checks)
  vector[4] phen_prob = [PrA, PrB, PrAB, PrO]';
п
set.seed(8878)
writeLines(stan_abo_missing_ab, con = "abo_missing_ab.stan")
mod_abo <- cmdstan_model("abo_missing_ab.stan")</pre>
fit_abo <- mod_abo$sample(</pre>
  data = list(n_A = 725, n_B = 258, n_0 = 1073, alpha = c(1,1,1)),
  seed = 8878, chains = 4, parallel_chains = 4,
  iter_warmup = 1000, iter_sampling = 1000
)
```